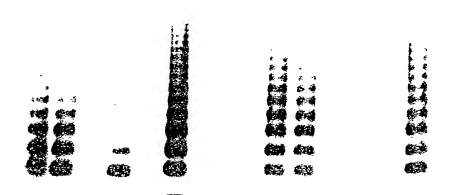
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Fig. 1: vWF Multimer Analysis Before and After
Anion Exchange Chromatography

A:+CaCl₂

B: -CaCl₂

A B a b c d e f



a: dissolved cryoprecipitate

b: Alu-supernatant

c: not bound to anion exchanger

d: 180 mM NaCl eluate +/- 10 mM CaCl₂

e: 200 mM NaCl eluate

f: 400 mM NaCl eluate

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Fig. 2: Detection of Factor II in Individual Fractions
Before and After Anion Exchange Chromatography

A B C D E F G



A: Factor II standard

B: dissolved cryoprecipitate

C: Alu-supernatant

The state was the state of the

D: 180 mM NaCl eluate

E: 400 mM NaCl eluate

F: 180 mM NaCl/+10 mM CaCl, eluate

G: 400 mM NaCl eluate

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Fig. 3: Protein S in the Individual Fractions

Before and After Anion Exchange Chromatography

ABCDEFG

 $PS_1 - PS_2 - PS_2$

A: Protein S standard

B: dissolved cryoprecipitate

C: Alu-supernatant

D: 180 mM NaCl eluate

E: 400 mM NaCl eluate

F: 180 mM NaCl/+10 mM CaCl, eluate

G: 400 mM NaCl eluate

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Fig. 4: Factor IX in the Individual Fractions

Before and After Anion Exchange Chromatography

A B C D E

-1

FIX —



A: Factor IX standard

B: dissolved cryoprecipitate

C: Alu-supernatant

D: 180 mM NaCl/10 mM CaCl₂ eluate

E: 400 mM NaCl eluate

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Fig. 5: Plasminogen in Individual Fractions
Before and After Anion Exchange Chromatography

A B C D

PG -- -- "

A: Plasminogen standard

B: dissolved cryoprecipitate

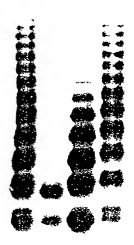
C: 400 mM eluate anion exchanger

D: eluate lysine-Sepharose

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Fig. 6: vWF-Multimer Analysis Before and After Heparin Affinity Chromatography

A B C D



A: Starting material before heparin affinity chromatography,

B: Factor VIII/vWF-complex eluate 160 mM NaCl,

C: Factor VIII/vWF-complex eluate 230 mM NaCl,

D: Factor VIII/vWF-complex eluate 300 mM NaCl

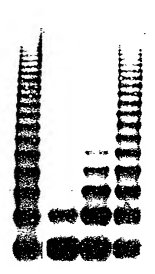
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Fig. 7: vWF Multimer Analysis of p-vWF and r-vWF Before and After Heparin Affinity Chromatography

A B C D

A B C D





I. p-vWF

II. r-vWF

A: p-vWF-starting material

A: r-vWF starting material

B: p-vWF/LMW

B: r-vWF/LMW

C: p-vWF/MMW

C: r-vWF/MMW

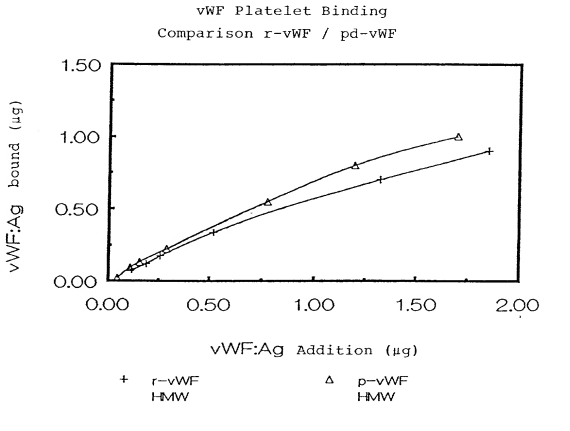
D: p-vWF/HMW

D: r-vWF/HMW

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Fig. 8 Comparison of the Binding of r-vWF/HMW and p-vWF/HMW to Platelets

Graphical representation of the amount of vWF added and of the platelet-bound amount of vWF



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Fig. 9 Binding of p-vWF/HMW and r-vWF/HMW to Platelets and Multimer Analysis

A: p-vWF/HMW;

B: r-vWF/HMW;

a: vWF, not bound;

b: platelet-bound vWF

c: vWF starting fraction after affinity chromatography

